VANMA48.001CP1 **PATENT**

ANTIGENIC POLYPEPTIDE SEQUENCES OF FACTOR VIII, AND FRAGMENTS AND/OR EPITOPES OF THESE SEQUENCES

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Cross-Reference to Related Applications

This application is a Continuation-in-Part of U.S. Application serial number 08/765,837, filed September 7, 1999, which was a U.S. National Phase application of PCT/BE95/00068, filed July 14, 1995, which claims priority to Belgian Application BE 9400666, filed July 14, 1994, the disclosures of which are incorporated herein by reference in their entireties.

Field of the Invention

The present invention relates to the antigenic polypeptide sequences of factor VIII, to fragments and epitopes of these sequences and to the major parts of these epitopes, to the inhibitors which are directed against these sequences, its fragments, its epitopes and/or major parts of these epitopes, and to anti-inhibitors which are directed against the said inhibitors.

The present invention also relates to a pharmaceutical composition and to a diagnostic device comprising at least one of the above mentioned molecules.

Background of the Invention

FVIII is a large multi-domain protein of 2,332 amino acids made up of three domains, A, B and C which are arranged in the order A1:a1:A2:a2:B:a3:A3:C1:C2. The A domains possess more than 40% homology and are also homologous to ceruloplasmin (for recent review, see Pratt (2000) and Saenko (1999)). 30% homology also exists between the A domains of factor V and FVIII. The C domain occurs twice and is reported to be able to bind glyco-conjugates and phospholipids having a net negative charge. It exhibits homology with lectins which are able to bind to negatively charged phospholipids. The platelet attachment site has been located in this region (C2 domain) (Foster et al., (1990)).

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These antigenic determinants consist of fragments 351 - 365 (A1 domain - heavy chain), 713 - 740 (A2 domain), 1670 - 1684 (A3 domain - light chain) (NH₂ end of the light chain) or else 2303 - 2332 (C2 domain - light chain) (Foster C, (1990)), fragments 701 - 750, 1663 - 1689, 330 - 472, 1694 - 1782 (EP-0 202 853), 322 - 740 and 2170 - 2322.

The U.S. patent 5,744,446 describes an hybrid human/animal Factor VIII having a sequence of amino acids selected from the group of the A2 domain fragments 373-540, 373-508, 445-508, 484-508, 404-508, 489-508 and 484-489, with corresponding sequences of porcine or murine Factor VIII, said hybrid being used for the treatment of Factor VIII deficiencies.

The antibodies which recognize these various sites interfere, with the activation of FVIII, the binding of vWf, FIXa, FXa, APC or phospholipids. The specific antibody response to FVIII vary considerably among individuals, and epitopes for inhibitor antibodies have to be determined for all FVIII domains (see for recent review Scandella, 2000; Lollar, 2000).

Other antibodies, which do not inhibit standard activity tests in vitro, can exert an influence on the behavior of FVIII with the other constituents of the coagulation cascade while attaching themselves to sites in the molecule which are at a substantial distance from the active sites. These antibodies, can interfere with the natural state of folding of FVIII by altering some of its properties.

Emergence of alloantibodies (inhibitors) that neutralize infused FVIII activity may seriously complicate FVIII replacement therapy. Reported inhibitor incidence rates in hemophiliacs vary considerably. They range around 6-35% (Vermylen et al, 1998). Candidates for genetic predispositions such as large deletions and intron 22 inversion have been found associated with a high incidence of inhibitors and genes that are involved in the immune response as genes MHC class I and class II (Tuddenham and McVey, 1998). Repeat switching from one FVIII product to another and the possibility that some FVIII concentrates are more immunogenic may also explain the appearance of inhibitors (Vermylen et al, 1998). Different methods of preparing FVIII could exert an influence on its structure, its physicochemical properties or its natural microenvironment; Laub et al. (1999); Raut et al. (1998)). Clinically relevant anti-FVIII

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autoantibodies are rare in non-hemophilic patients (annual frequency in the population: 1-5/10⁶) (Morrisson and Ludlam) (1995). They are associated with a number of autoimmune diseases and are often characterized by life-threatening hemorrhage. On the other hand, anti-FVIII antibodies have also been described in healthy subjects (Algiman et al, 1992; Moreau et al, 2000), without any apparent effect on the subjects' levels of circulating FVIII.

Self proteins or derived peptides may elicit an immune response if presented to CD4 T cells at inflammatory sites by professional antigen presenting cells. Using pools of overlapping synthetic peptides spanning the sequences of individual FVIII domains, Reding et al. (2000) showed reactive CD4⁺ to FVIII in healthy subjects and hemophilia patients. Several FVIII domains were recognized: A3 domain was recognized more strongly and frequently and each domain forms several epitopes.

Techniques such as western blotting, immunoprecipitation, and enzyme-linked immunosorbent assays (ELISAs), using well-defined FVIII proteolytic fragments, a large recombinant peptide library, or synthetic peptide arrays, have been used to map different FVIII-inhibitor binding sites located mainly in the A2 and C2 domains. However, none of these techniques has made it possible to build a model for identification of inhibitor and non-inhibitor epitopes. Only a few epitopes have been mapped to discrete sequences (<20 amino-acid residues). To solve this problem, Palmer et al (1997) synthesized 96 undecamer peptides (11 amino-acid residues) representing 80% of the complete residue sequence of FVIII. They succeeded in determining the epitope specificity of 9 patients' inhibitory antibodies. Other useful techniques are analysis of FVIII gene mutations and their effects on the FVIII molecule as well as phage display technology (van den Brink et al, 2000). All these methodologies, however, are time consuming, rather costly, and largely dependent on patient availability. Certain areas of the FVIII molecule may be "hot spots" containing commonly recognized clusters of inhibitor epitopes, e.g., regions in the A2 domain, A3 domain, and C2 domain. The reason for these "hot spots" in generating an inhibitor response remains poorly understood (Reisner et al, 1995).

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Currently, a predominant notion among hemophilic patients, clinicians and "fractionators" is that of having available a purified FVIII which is devoid of all pathogenic plasma contaminants and secondary effects.

Different animal models could be used as hemophilia dogs, SCID mice, hemophilia mice ... but until now, no satisfactory experimental model exists which makes it possible to forecast the immunogenicity or the immuno-modulatory effect of the FVIII preparations, or the susceptibility of the host, before they have been administered clinically.

Patients who develop an anti-FVIII immune response find themselves in a serious situation which necessitates the use of severe, aggressive and excessively expensive measures.

One of the frequently treatment, is the induction of immune tolerance by administration of very high doses of FVIII (150 IU/kg twice a day) in association or not with prothrombin complex concentrates and is assigned as "Bonn Protocol". Treatment options are also to by-pass the FVIII inhibitor activity by use of PCC (preferably an activated PCC [APCC]) or FVIIa. Specific antibodies as consequence of the infusion of these alternative agents could be produced, impairing the treatment. As an alternative agent porcine FVIII may be used to achieve haemostasis in patients with antibodies that do not substantially crossreact with porcine FVIII before or during the treatment (Lollar, 2000).

A potential alternative approach to inhibit the production of inhibitors is blockade of the T cell/B cell collaboration mediated by through receptor ligand binding signal events (Ewenstein et al, 2000). Preliminary clinical trials were performed using a humanized mouse monoclonal antibody to human T cell CD40 ligand (CD 154).

A profitable strategy for reducing the level of inhibitors has consisted in subjecting patients to an extracorporeal circulation to enable solid-phase absorption of the total IgG.

The immunoabsorbant could be sepharose-bound staphylococcal protein A or sepharose-bound polyclonal sheep antibodies to total human immunoglobulin (Knobf and Derfler, 1999). The foreign proteins (protein A, sheep anti-human Ig) could leak

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from the column and triggered the immune system of the recipient; moreover problems could raised as sanitization (ICH Topic Q5A, Directive 92/79/EC).

The infusion of polyvalent intravenous immunoglobulins (IVIG), where appropriate combined with an immunosuppressive treatment, has been found to be relatively effective, although the reason for this effectiveness is still not fully established. Various hypotheses involving feed-back inhibition of IgG synthesis, stimulation of IgG clearance or activation of T suppressor cells have been advanced. An interesting explanation is that these commercial intravenous immunoglobulins might contain antibodies which are able to react with the variable parts (idiotypes) of the anti-FVIII antibodies and neutralize these antibodies (Dietrich et al. (1992)).

Unfortunately, none of these approaches has been found to be satisfactory in terms of safety, efficacy, efficiency and cost.

The state of the art in epitope structure prediction was limited given to the fact that non-continuous amino acid residues seem to constitute most important epitope and that the dynamics of binding is often not integrated into the epitope prediction equation making epitope structure prediction a complex four-dimensional problem (Van Regenmortel, Methods: A companion to Methods in Enzymology, 9, page 465-472, 1996).

According to the author, most of the antibodies raised against intact proteins do not react with any peptide fragment derived from the parent protein indicating that such antibodies are directed to discontinuous epitopes (conformational epitopes).

This author states also that low success rate of antigenic prediction is due to the fact that predictions concerns only continuous epitopes and it is unrealistic to reduce the complexity of epitopes that always possess conformational features to one dimensional linear peptide model.

Similarly, Palmer et al. (1997) using synthetic peptide arrays to identify novel Factor VIII inhibitor epitopes note that each patient pattern of anti-factor VIII antibody reactivity appears to be polyclonal, directed against multiple sites located within the amino and carboxyl terminus of the protein and seems to be unique for each plasma investigated (see also above).

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Moreover, this author notes that it is difficult to predict the importance that any given antibody: epitope interaction may have on Factor VIII coagulation activity based on the results of synthetic peptide assays alone (due to the incomplete understanding of the relationship between structure and function of different factor VIII domains and the possibility that both inhibitor and non-inhibitory antibodies may be present in a patient's plasma.

Therefore, the documents of the state of the art do not suggest to identify antigenic linear peptides upon a macro-molecule (such as Factor VIII) and that linear epitopes could be used for the diagnostic and/or the therapy of immune disorders induced by inhibitors directed against Factor VIII.

The present invention aims to obtain antigenic polypeptide sequences of factor VIII, fragments and epitopes of these sequences, whose purpose is to improve the diagnosis and/or therapy (including prevention) of immune disorders (in particular those induced by inhibitors of FVIII and inhibitors of FVIII, especially inhibitors of the binding of the von Willebrand factor (vWf), to the FIX and/or to membrane phospholipids (PL)), and which allows a screening between non-inhibitory and inhibitory anti-FVIII allo- or auto-antibodies (allo- or auto-immunoglobulins).

Another aim of the invention is to obtain inhibitors which exhibit an immunoaffinity with these antigenic polypeptide sequences, fragments and/or epitopes, as well as to obtain anti-inhibitors, in particular antibodies or (T)cell receptors, which are directed against the above-mentioned said inhibitors and whose purpose is to improve the diagnosis and/or therapy (or prevention) of immune disorders.

A further aim of the invention is to obtain said molecules at high purity, in industrial level, without contaminants (viruses, prions,...) and according to the GMP practices in the field of therapy and diagnostics (ICH topic QSA, Directive 92/79/EC, etc.).

Summary of the Invention

Some embodiments of the present invention are described in the following numbered paragraphs.

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Paragraph 1: An antigenic polypeptide sequence, which is the polypeptide sequence of factor VIII.

Paragraph 2: An antigenic polypeptide sequence, which lacks the following fragments: alanine 322 - serine 750, leucine 1655 - arginine 1689, lysine 1694 - proline 1782 and possibly the fragment aspartic acid 2170 - tyrosine 2332.

Paragraph 3: The sequence according to paragraph 1 or 2, which is immunogenic.

Paragraph 4: The sequence according to paragraph 3, which exhibits an immunoaffinity for the receptors of T and/or B lymphocytes.

Paragraph 5: An antigenic fragment of the sequence according to paragraph 1 or 2, which is selected from the group consisting of the polypeptide sequences A1, A2, A3 or C of factor VIII.

Paragraph 6: The antigenic fragment of the polypeptide sequence A3 according to paragraph 5, which is selected from the group consisting of the sequence fragment arginine 1649 to arginine 2031 inclusive, the sequence fragment threonine 1739 to aspartic acid 1831 inclusive and/or the sequence fragment arginine 1803 to arginine 1917 inclusive.

Paragraph 7: A sequence epitope of the fragment according to paragraph 6, which is selected from the group consisting of:

the epitope arginine 1648 to tyrosine 1664 inclusive, defined by the following sequence:

SEQ ID NQ:1:

Arg Asp Ile Thr Arg Thr Thr Leu Gln Ser Asp Gln Glu Glu Ile Asp Tyr,

possibly deleted from one or more amino acids of the tetrapeptide Arg-Asp-Ile-Thr or one or two of the last amino acids of the peptide Asp-Tyr,

the epitope aspartic axid 1681 to arginine 1696 inclusive, defined by the following sequence:

SEQ ID NO:2:

Asp Glu Asp Glu Asn Gln Ser Pro Arg Ser Phe Gln Lys Lys Thr Arg, possibly deleted from one or more amino acids of the epitope Asp-Glu-Asp-Glu,



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the epitope threonine 1739 to tyrosine 1748 inclusive, defined by the following sequence:

SEQ ID NO:3:

Thr Asp Gly Ser Phe Thr Gln Pro Leu Tyr,

the epitope\asparagine 1777 to phenylalanine 1785 inclusive, defined by the following sequence:

SEQ ID NO:4:

Asn Gln Ala Ser Arg Pro Tyr Ser Phe,

possibly deleted from one or two amino acids of the terminal dipeptide Ser-Phe or the tetrapeptide Pro-Tyr-Ser-Phe,

the epitope glutamic acid 1794 to tyrosine 1815 inclusive, defined by the following sequence:

SEQ ID NO:5:

Glu Asp Gln Arg Gln Gly Ala Glu Pro Arg Lys Asn Phe Val Lys Pro Asn Glu Thr Lys Thr Tyr,

possibly deleted from one or more amino acids from the first tripeptide Glu-Asp-Gln or the first nonapeptide Glu-Asp-Gln-Arg-Gln-Gly-Ala-Glu-Pro,

the epitope methionine 1823 to aspartic acid 1831, defined by the following sequence:

SEQ ID NO:6:

Met Ala Pro Thr Lys Asp Glu Phe Asp,

the epitope glutamic acid 1885 to phenylalanine 1891 inclusive, defined by the following sequence:

SEQ ID NO:7:

Glu Thr Lys Ser Trp Tyr Phe,

the epitope glutamic acid 1885 to alanine 1901 inclusive, defined by the following sequence:

SEQ ID NO:8:

Glu Thr Lys Ser Trp Phe Thr Glu Asn Met Glu Arg Asn Cys Arg Ala,

possibly deleted from one or more amino acids from the heptapeptide Gly-Thr-Lys-Ser-Trp-Phe-Thr or from the tripeptide Cys-Arg-Ala,

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the epitope aspartic acid 1909 to arginine 1917 inclusive, defined by the following sequence: SEQ ID NO:9: Asp\Pro Thr Phe Lys Glu Asn Tyr Arg, and the epitope comprised between serine 2018 and histidine 2031 inclusive, defined by the following sequence: SEQ ID NO:10: Ser Asn Lys Cys Gln Thr Pro Leu Gly Met Ala Ser Gly His. Paragraph 8: An antigenic fragment of the polypeptide sequence A1 according to paragraph 5, which is alanine 108 to methionine 355 inclusive, preferably alanine 108 to alanine 227 inclusive. Paragraph 9: A sequence epitope of the fragment according to paragraph 8, which is selected from the group consisting of: the epitope alanine 108 to valine 128 inclusive, defined by the following sequence: SEQ ID NO:11: Ala Ser Glu Gly Ala Glu Tyr Asp Asp Gln Thr Ser Gln Arg Glu Lys Glu Asp Asp Lys Val, possibly deleted from the terminal amino acids alanine and/or valine, the epitope glutamic acid 181 to leucine 192 inclusive, defined by the following sequence: SEO ID NO:12: A possibly deleted from one or two amino acids of the terminal dipeptide Thr-Leu, the epitope aspartic acid 203 to alanine 227 inclusive, defined by the following sequence: SEQ ID NO:13:

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Asp Glu Gly Lys Ser Trp His Ser Glu Thr Lys Asn\Ser Leu Met Gln Asp Arg Asp Ala Ala Ser

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Ala Arg Ala,

possibly deleted from one or more amino acids of the nonapeptide Asp-Arg-Asp-Ala-Ala-Ser-Ala-Arg-Ala,

and the epitope aspartic acid 327 to methionine 355 inclusive, defined by the following sequence:

SEQ 10 NO:14:

Asp Set Cys Pro Glu Glu Pro Gln Leu Arg Met Lys Asn Asn Glu Glu Ala Glu Asp Tyr Asp Asp Asp Ser Glu Met,

possibly deleted from one or more amino acids of the dipeptide Asp-Ser or the octapeptide Asp-Asp-Leu-Thr-Asp-Ser-Glu-Met,

Paragraph 10: An antigenic fragment of the antigenic polypeptide sequence A2 according to paragraph 5, which is aspartic acid 403 to aspartic acid 725 inclusive, preferably histidine 693 to aspartic acid 725 inclusive.

Paragraph 11: A sequence epitope of the fragment according to paragraph 10, which is selected from the group consisting of:

the epitope aspartic acid 403 to lysine 425 inclusive, defined by the following sequence:

SEQ ID NO:15:

Asp Asp Arg Ser Tyr Lys Ser Gln Tyr Leu Asn Asn Gly Pro Gln Arg Ile Gly Arg Lys

Tyr Lys Lys.

possibly deleted from one or more amino acids of the tetrapeptide Asp-Asp-Arg-Ser,

the epitope valine 517 to arginine 527 inclusive, defined by the following sequence:

SEQ ID NO:16:

Val Glu Asp Gly Pro Thr Lys Ser Asp Pro Arg,

possibly deleted from one or the two amino acids of the dipeptide Pro-Arg,

the epitope tyrosine 555 to glutamine 565 inclusive defined by the following sequence:

SEQ ID NO:17:

Tyr Lys Glu Ser Val Asp Gly Arg Gly Asn Gln,

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the epitope histidine 693 to glycine 701 inclusive, defined by the following sequence

SEO ID NO:18

His Asn Ser Asp Phe Arg Asn Arg Gly,

the ephtope serine 710 to aspartic acid 725 inclusive, defined by the following sequence:

SEQ ID NO:19

Ser Cys Asp Lys Asn Thr Gly Asp Tyr Try Gly Asp Ser Tyr Glu Asp,

the epitope leucine 730 to serine 741 inclusive, defined by the following sequence:

SEQ ID NO:20:

Leu Leu Ser Lys Asn Ala Ile Glu Pro Arg Ser,

possibly deleted from the terminal amino acid serine and/or the first amino acid leucine,

the epitope serine 817 to serine 830 inclusive, defined by the following sequence:

SEQ ID NO:21:

Ser Asp Asp Pro Ser Gly Ala Ile Asp Ser Asn Asn Ser.

Paragraph 12: An antigenic fragment of the antigenic polypeptide sequence C according to paragraph 5, which is lysine 2085 to isoleucine 2251 inclusive, or leucine 2273 to tyrosine 2332 inclusive, preferably lysine 2085 to glycine 2121 inclusive or serine 2182 to leucine 2251 inclusive.

Paragraph 13: A sequence epitope of the fragment according to paragraph 12, which is selected from the group consisting of:

the epitope isoleucine 2081 to serine 2095 inclusive, defined by the following sequence

SEQ ID NO:22:

Ile His Gly Ile Lys Thr Gln Gly Ala Arg Gln Lys Phe Ser Ser,

possibly deleted from one or more amino acids of the texapeptide Ile-His-Gly-

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the epitope tyrosine 2105 to glycine 2121 inclusive, defined by the following sequence:

SEQ ID NO:23:

Tyr Ser Leu Asp Gly Lys Lys Trp Gln Thr Tyr Arg Gly Asn Ser Thr Gly, possibly deleted from one or more amino acids of the tripeptide Tyr-Ser-Leu, the epitope asparagine acid 2128 to asparagine acid 2138 inclusive, defined by the following sequence:

SEQ ID NO:24:

Asn Val Asp Ser Ser Gly Ile Lys His Asn,

the epitope histidine 2152 to arginine 2163 inclusive, defined by the following sequence:

SEQ ID NO:25:

His Pro Thr His Tyr Ser Ile Arg Ser Thr Leu Arg,

the epitope serine 2181 to asparagine acid 2198 inclusive, defined by the following sequence:

SEQ ID NO:26:

Ser Lys Ala Ile Ser Asp Ala Gln Ile Thr Ala Ser Ser Tyr Phe Thr Asn,

possibly deleted from one or more amino acids of the first dipeptide Ser-Tyr or one or more amino acids from the terminal tripeptide Phe-Thr-Asn,

the epitope serine 2204 to glutamine 2222 inclusive, defined by the following sequence:

SEQ ID NO:27:

Ser Pro Ser Lys Ala Arg Leu His Leu Gln Gly Arg Ser Asn Ala Trp Arg Pro Gln,

the epitope glutamine 2235 to leucine 2251 inclusive, defined by the following sequence:

SEQ ID NO:28:

Gln Lys Thr Met Lys Val Thr Gly Val Thr Thr Gln Gly Val Lys Ser Leu,

possibly deleted from one or two amino acids of the terminal dipeptide Ser-Leu or one or more amino acids of the tetrapeptide Val-Lys-Ser-Leu,

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the epitope glycine 2242 to leucine 2251 inclusive, defined by the following sequence:

SEQ ID NO:29:

Gly Val Thr Thr Gln Gly Val Lys Ser Leu,

possibly deleted from one or two amino acids of the terminal dipeptide Ser-Leu, the epitope isoleucine 2262 to glutamine 2270 inclusive, defined by the following sequence:

SEQ ID NO:30:

Ile Ser Ser Ser Gln Asp Gly His Gln,

the epitope leucine 2273 to serine 2289 inclusive, defined by the following sequence:

SEQ ID NO:31:

Leu Phe Phe Gln Asn Gly Lys Val Lys Val Phe Gln Gly Asn Gln Asp Ser,

the epitope proline 2292 to tyrosine 2305 inclusive, defined by the following sequence:

SEQ ID NO:32:

Pro Val Val Asn Ser Leu Asp Pro Pro Leu Leu Thr Arg Tyr,

possibly deleted from one or more amino acids of the terminal tripeptide Thr-Arg-Tyr,

the epitope glutamic acid 2322 to tyrosine 2332 inclusive, defined by the following sequence:

SEQ ID NO:33:

Glu Val Leu Gly Cys Glu Ala Gln Asp Leu Tyr.

Paragraph 14: A conformational epitope, which contains at least two different epitopes according to any one of the preceding paragraphs 8, 10 and 12.

Paragraph 15: A pool of more than three fragments or epitopes according to any of the preceding paragraphs 6 to 14.

Paragraph 16: A recombinant factor VIII having an amino acid sequence deleted from one or more of the fragments or the epitopes according to any one of the paragraphs 6 to 14.

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Paragraph 17: A complex comprising a carrier protein or a carrier peptide linked to an element which is selected from the group consisting of the fragment and/or the epitope according to any one of the paragraphs 6 to 14.

Paragraph 18: An inhibitor of factor VIII, which exhibits an immunoaffinity with the sequence, the fragment, the epitope, the pool and/or the complex according to any one of the preceding paragraphs.

Paragraph 19: The inhibitor according to paragraph 18, which is an anti-factor VIII antibody or antibody fragment.

Paragraph 20: An anti-inhibitor, which is directed against the inhibitor of factor VIII according to paragraph 18 or 19.

Paragraph 21: The anti-inhibitor according to paragraph 20, which is an anti-anti-factor VIII idiotype antibody or antibody fragment.

Paragraph 22: A pharmaceutical composition, which comprises an adequate pharmaceutical carrier and at least one element selected from the group consisting of the sequence, the fragment, the epitope, the pool, the complex, the recombinant factor VIII or the inhibitor and/or the anti-inhibitor according to any one of the preceding paragraphs.

Paragraph 23: A diagnostic and/or purification device, which comprises at least one element which is selected from the group consisting of the sequence, the fragment, the epitope, the pool, the complex, the inhibitor and/or the anti-inhibitor according to any one of the preceding paragraphs.

Paragraph 24: The device according to paragraph 23, which is a diagnostic kit.

Paragraph 25: The device according to paragraph 23, which is a chromatography column or filter.

Paragraph 26: A method for a therapeutic treatment and/or prevention of an immune disorder in mammal, wherein the pharmaceutical composition according to paragraph 22 is administered to the mammal patient presently or potentially having said immune disorder, in an amount effective to treat and/or prevent said immune disorder.

Paragraph 27: A method for a therapeutic treatment and/or prevention of an immune disorder in a mammal patient, wherein a physiological fluid such as serum obtained from said mammal patient is put into the chromatography column of paragraph

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25 in order to allow a binding with the inhibitors of factor VIII present in said serum with the sequence, the fragment, the epitope, the pool and/or the complex according to any of the preceding paragraphs 1 to 15, wherein the physiological liquid is eluted from said chromatography column and the physiological liquid from which the inhibitors of factor VIII have been removed is reinjected to the patient.

Paragraph 28: The therapeutic treatment and/or prevention method according to paragraph 26 or 27, wherein the immune disorder is induced by an element selected from the group consisting of inhibitors of factor VIII, inhibitors of the binding of factor VIII to the von Willebrand factor, to the factor IX, the factor X and/or to membrane phospholipids.

Paragraph 29: A process for identifying and obtaining inhibitors and/or antiinhibitors according to paragraph 18 or 19, comprising the steps of:

selecting an element from the group consisting of the sequence, the fragment, the epitope, the pool and/or the complex according to any one of the preceding paragraphs 1 to 15 or 17, attached to a solid support of a chromatography column,

passing a physiological fluid from a patient containing inhibitors of factor VIII through said chromatography column,

eluting said column, and

collecting the fractions containing inhibitors of factor VIII which have exhibited an immunoaffinity with said element.

Paragraph 30: The process according to paragraph 29, further comprising the steps of:

attaching the collected inhibitors of factor VIII upon the solid support of a chromatography column,

passing a physiological fluid from a patient containing anti-inhibitors of factor VIII through said chromatography column,

eluting said column, and

collecting the fractions containing anti-inhibitors of factor VIII which have exhibited an immunoaffinity with said inhibitors of factor VIII.

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Brief Description of the Drawings

Figure 1. This figure depicts the hydrophilicity, flexibility and accessibility graph of the A3 sequence of Factor VIII, renumbered 1 to 371 amino acids (surface value for each amino acid).

Figure 2. This figure (2a) represents the elution profile related to the purification of human anti-SEQ. ID NO: 32 antibodies by affinity chromatography in peptide-Sepharose column. Cohn fraction II+III solution (50 mL) was loaded onto the column (1 mL gel) at a flow rate of 1 mL/min. The separation of specific antibodies was performed as described in the Examples. The arrow indicates the position of specific human anti-SEQ. ID NO: 32. IgG purified from Cohn fraction II+III. The clotting activity of FVIII (2b) was measured as described in Examples in the presence of increasing amount of anti-SEQ. ID NO: 32. The of FVIII activity = (FVIII activity in the presence of antibody/FVIII activity in the absence of antibody) *100.

Figure 3. This figure represents the human anti-peptide antibody immunoreactions with FVIII polypeptides after Western Blotting (panel A from left to right: human antibodies HAP1 through HAP4, specific for different FVIII epitope sequences found in the FVIII HC -- see also Table 2 and panel B: human antibodies specific for the P5 peptide and the FVIII LC sequences, P7, P8, and P9 --see also Table 2). The RAP9 lane shows the reactivity of FVIII polypeptieds towards purified rabbit antibodies specific for the peptide sequence Arg¹⁷⁹⁷–Tyr¹⁸¹⁵ (see also Table 2).

Figure 4. This figure represents the ELISA reactivity of the four inhibitor plasmas with different peptide sequences. Inhibitors present in four patient plasmas were analyzed by ELISA test using as coated antigens the different selected FVIII epitopes synthetic peptides, as indicated by the ordinate.

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Detailed Description of the Preferred Embodiment

The present invention relates to the antigenic polypeptide sequences of factor VIII and/or fragments of these sequences, as described by Verhar et al. (1984), the disclosure of which is incorporated herein by reference in its entirety.

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The "polypeptide sequence of factor VIII" is understood to be the natural human or animal sequence, which may be glycosylated and which has been obtained by

purification from pools of plasma, in particular cryoprecipitate, by synthesis and/or by genetic manipulation (sequence from which portions which are not involved in the mechanism of blood coagulation may have been deleted) of factor VIII.

The present invention relates, in particular, to the antigenic polypeptide sequences of factor VIII which lacks the fragments comprised between alanine 322 - serine 750, leucine 1655 - arginine 1689 and lysine 1694 - proline 1782, and possibly also the fragments comprised between aspartic acid 2170 and tyrosine 2332.

The present invention relates, in particular, to the antigenic polypeptide sequences A1, A2, A3 and C (C1 and C2) of factor VIII.

A first embodiment of the invention relates to the antigenic polypeptide sequence A3 of factor VIII, and to fragments and/or epitopes of this sequence. The said sequence contains the fragments glutamic acid 1649 to histidine 2031 inclusive, arginine 1652 to arginine 1917 inclusive or arginine 1803 to arginine 1917 inclusive, of the polypeptide sequence of factor VIII as published by Verhar et al. (1984) and Toole et al. (1984).

Preferably, the fragments of the said sequence are arginine 1648 to arginine 1696 inclusive, threonine 1739 to aspartic acid 1831 inclusive or glutamic acid 1885 to arginine 1917 inclusive.

The fragments, epitopes and major parts thereof are preferably polypeptidic sequences made of at least 7 amino acids of the FVIII polypeptidic sequence.

The invention also relates to the sequence epitopes of these fragments, in particular:

- the epitope arginine 1648 to tyrosine 1664 inclusive, defined by the following sequence:

SEQ ID NO.:1:

Arg Asp Ile Thr Arg Thr Thr Leu Gln Ser Asp Gln Glu Glu Ile Asp Tyr, and possibly deleted from one or more amino acids of the tetrapeptide Arg-Asp-Ile-Thr (P7), or one or two of the last amino acids of the dipeptide Asp-Tyr

- the epitope aspartic acid 1681 to arginine 1696 (P8) inclusive, defined by the following sequence:

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SEQ ID NO.:2:

Asp Glu Asp Glu Asn Gln Ser Pro Arg Ser Phe Gln Lys Lys Thr Arg

possibly deleted from one or more amino acids of the epitope Asp-Glu-Asp-Glu,

5 - the epitope threonine 1739 to tyrosine 1748 inclusive, defined by the following sequence:

SEQ ID NO.:3:

Thr Asp Gly Ser Phe Thr Gln Pro Leu Tyr

- the epitope asparagine 1777 to phenylalanine 1785 inclusive, defined by the following sequence:

SEQ ID NO.:4:

Asn Gln Ala Ser Arg Pro Tyr Ser Phe

possibly deleted from one or more amino acids of the terminal dipeptide Ser-Phe or tetrapeptide Pro-Tyr-Ser-Phe

the epitope glutamic acid 1794 to tyrosine 1815 inclusive, defined by the following sequence:

SEQ ID NO.:5:

Glu Asp Gln Arg Gln Gly Ala Glu Pro Arg Lys Asn Phe Val Lys Pro

Asn Glu Thr Lys Thr Tyr,

possibly deleted from one or more amino acids of the first tripeptide Glu-Asp-Gln (P9) or the first nonapeptide Glu-Asp-Gln-Arg-Gln-Gly-Ala-Glu-Pro

- the epitope methionine 1823 to aspartic acid 1831 inclusive, defined by the following sequence:

SEQ ID NO.:6:

Met Ala Pro Thr Lys Asp Glu Phe Asp

- the epitope glutamic acid 1885 to phenylalanine 1891 inclusive, defined by the following sequence:

SEQ ID NO.:7:

Glu Thr Lys Ser Trp Tyr Phe

- the epitope glutamic acid 1885 to alanine 1901 inclusive, defined by the following sequence:

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SEQ ID NO.:8:

Glu Thr Lys Ser Trp Phe Thr Glu Asn Met Glu Arg Asn Cys Arg Ala possibly deleted from one or more amino acids from the heptapeptide Glu-Thr-Lys-Ser-Trp-Phe-Thr or from the tripeptide Cys-Arg-Ala.

- the epitope aspartic acid 1909 to arginine 1917 inclusive, defined by the following sequence:

SEQ ID NO.:3:

Asp Pro Thr Pha Lys Glu Asn Tyr Arg

- the epitope comprised between serine 2018 and histidine 2031 inclusive, defined by the following sequence:

SEQ ID NO.:10:

Ser Asn Lys Cys Gln Th\Pro Leu Gly Met Ala Ser Gly His

Advantageously, the said sequences, specific fragments and epitopes exhibit an antigenic characteristic which is iNustrated by Table 1.

Another preferred embodiment of the invention relates to antigenic polypeptide sequence A1 of factor VIII, fragments and/or epitopes of this sequence.

Preferably, the fragments of the said sequence are alanine 108 to methionine 355 inclusive, preferably alanine 108 to alanine 227 inclusive.

The invention also relates to the sequence epitopes of these fragments, in particular:

- the epitope alanine 108 to valine 128 inclusive, defined by the following sequence:

SEQ ID NO.:11:

Ala Ser Glu Gly Ala Glu Tyr Asp Asp Gln Thr Ser Gln Arg Glu Lys

Glu Asp Asp Lys Val

possibly deleted from the terminal amino acids alanine and valine (P1)

- the epitope glutamic acid 181 to leucine 192 inclusive, defined by the following sequence:

SEQ ID NO.:12:

Glu Gly Ser Leu Ala Lys Glu Lys Thr Gln Thr Leu

possibly deleted from one or two amino acids of the terminal dipertide Thr-Leu

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- the epitope aspartic acid 203 to alanine 227 inclusive, defined by the following sequence:

SEQ ID NO.:13:

Asp Glu Gly Lys Ser Trp His Ser Glu Thr Lys Asn Ser Leu Met Gln

Asp Arg Asp Ala Ala Ser Ala Arg Ala

possibly deleted from one or more amino acids of the nonapeptide Asp-Arg-Asp-Ala-Ala-Ser-Ala-Arg-Ala

- the epitope aspartic acid 327 to methionine 355 inclusive, defined by the following sequence:

10 SEQ ID NO.:\\4:

Asp Ser Cys Pro Glu Glu Pro Gln Leu Arg Met Lys Asn Asn Glu Glu

Ala Glu Asp Tyr Asp Asp Asp Leu Thr Asp Ser Glu Met

possibly deleted from one or more amino acids from the terminal dipeptide Asp-Ser or the octapeptide Asp-Asp-Leu-Thr-Asp-Ser-Glu-Met (P2).

Another preferred embodiment of the invention relates to the antigenic polypeptide sequence A2 of factor VIII, fragments and/or epitopes of this sequence.

Preferably, the fragments of the said sequence are aspartic acid 403 to serine 840 inclusive, preferably histidine 693 to aspartic acid 725 inclusive.

The invention also relates to the sequence epitopes of these fragments, in particular:

the epitope aspartic acid 403 to lysine 425 inclusive, defined by the following sequence:

SEQ ID NO.:15:

Asp Asp Arg Ser Tyr Lys Ser Gln Tyr Leu Asn Asn Gly Pro Gln Arg

Ile Gly Arg Lys Tyr Lys Lys
possibly deleted from one or more amino acids of the tetrapeptide Asp-Asp-ArgSer (P3),

the epitope valine 517 to arginine 527 inclusive, defined by the following sequence: SEQ ID NO.:16:

Val Glu Asp Gly Pro Thr Lys Ser Asp Pro Arg
possibly deleted from one or the two amino acids of the dipeptide Pro-Arg,

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-\ the epitope tyrosine 555 to glutamine 565 inclusive, defined by the following sequence:

SEQ ID NO.:17:

Tyr Lys Glu Ser Val Asp Gly Arg Gly Asn Gln

- the epitope histidine 693 to glycine 701 inclusive, defined by the following sequence:

SEQ ID NO.:18:

His Asn Sex Asp Phe Arg Asn Arg Gly

- the epitope serine 710 to aspartic acid 725 inclusive, defined by the following sequence (P4):

SEQ ID NO.:19:

Ser Cys-Asp Lys Ash Thr Gly Asp Tyr Try Gly Asp Ser Tyr Glu Asp

the epitope leucine 730 to serine 741 inclusive, defined by the following sequence (P4):

SEQ ID NO.:20:

Leu Leu Ser Lys Asn Asn Ala Ile Glu Pro Arg Ser

possibly deleted from the terminal amino acid serine (P4) and/or the first amino acid leucine

the epitope serine 817 to serine 830 inclusive, defined by the following sequence (P5):

SEQ ID NO.:21:

Ser Asp Asp Pro Ser Gly Ala Ile Asp Ser Asn Asn Ser

A final preferred embodiment of the invention relates to the antigenic polypeptide sequence C of factor VIII, and fragments and/or epitopes of this sequence. Preferably, the fragments of the said sequence are histidine 2082 to lysine 2251 inclusive or leucine 2273 to tyrosine 2332 inclusive, preferably lysine 2085 to glycine 2121 inclusive and serine 2181 to leucine 2251 inclusive.

The invention also relates to the sequence epitopes of these fragments, in particular:

- the epitope isoleucine 2081 to serine 2095 inclusive, defined by the following sequence:

SEQ ID NO.:22:

Ile His Gly Ile Lys Thr Gln Gly Ala Arg Gln Lys Phe Ser Ser possibly deleted from one or more amino acids from the tetrapeptide Ile-His-Gly-Ile

5 - the epitope tyrosine 2105 to glycine 2121 inclusive, defined by the following sequence:

SEQ ID NQ.:23:

Tyr Ser Leu Asp Gly Lys Lys Trp Gln Thr Tyr Arg Gly Asn Ser Thr

Gly

possibly deleted from one or more amino acids of the tripeptide Tyr-Ser-Leu (P10)

the epitope asparagine 2128 to asparagine 2138 inclusive, defined by the following sequence:

SEQ ID NO.:24:

Asn Val Asp Ser Ser Gly he Lys His Asn

the epitope histidine 2152 to arginine 2163 inclusive, defined by the following sequence:

SEQ ID NO.:25:

His Pro Thr His Tyr Ser Ile Arg Sen Thr Leu Arg

the epitope serine 2181 to asparagine 2198 inclusive, defined by the following sequence:

SEQ ID NO.:26:

Ser Lys Ala Ile Ser Asp Ala Gln Ile Thr Ala Ser Ser Tyr Phe Thr Asn possibly deleted from one or more amino acids from the terminal tripeptide Phe-Thr-Asn (P11)

- the epitope serine 2204 to glutamine 2222 inclusive, defined by the following sequence (P12):

SEQ ID NO.:27:

Ser Pro Ser Lys Ala Arg Leu His Leu Gln Gly Arg Ser Asn Ala Trp

30 Arg Pro Gln

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- the epitope glutamine 2235 to leucine 2251 inclusive, defined by the following sequence (P13):

SEQ ID NO.:28:

Gln Lys Thr Met Lys Val Thr Gly Val Thr Thr Gln Gly Val Lys Ser Leu possibly deleted from one or two amino acids of the terminal dipeptide Ser-Leu or one or more amino acids of the tetrapeptide Val-Lys-Ser-Leu

- the epitope glycine 2242 to leucine 2251 inclusive, defined by the following sequence:

SEQ ID NO.:29:

Gly Val Thr Thr Gln Gly Val Lys Ser Leu

possibly deleted from one or two amino acids of the terminal dipeptide Ser-Leu, said epitope presenting a possible partial overlapping with a known monoclonal antibody binding site ESH8 2248-2285

the epitope isoleucine 2262 to glutamine 2270 inclusive, defined by the following sequence:

SEQ ID NO.:30:

Ile Ser Ser Gln Asp Gly His Gln

the epitope leucine 2273 to serine 2289 inclusive, defined by the following sequence (P14):

SEQ ID NO.:31:

Leu Phe Phe Gln Asn Gly Lys Val Lys Val Phe Gln Gly Asn Gln Asp Ser

- the epitope proline 2292 to tyrosine 2305 inclusive, defined by the following sequence (P)(5):

SEQ ID/NO.:32:

Pro Val Val Asn Ser Leu Asp Pro Pro Leu Leu Thr Arg Tyr
possibly deleted from one or more amino acids of the terminal tripeptide

Thr-Arg-Tyr involved in the phospholipid von Willebrand factor binding site

- the epitope glutamic acid 2322 to tyrosine 2332 inclusive, defined by the following sequence (P16):

SEQ ID NO.:33:

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Glu Val Leu Gly Cys Glu Ala Gln Asp Leu Tyr

The invention also relates to the major parts of the said epitopes or the said fragments. Said epitopes can be deleted from one or more terminal amino acids, preferably from one, two or three amino acids, or can be replaced by one or more amino acids that present the same characteristic of hydrophilicity, flexibility and accessibility.

It is also known that some of the epitopes according to the invention are comprised in major determinants of human inhibitors epitopes or several factors binding sites or binding sites of known monoclonal antibodies, especially the portion C2 that is known to be the binding site of the monoclonal antibody Mas531P or the binding site ESH8 as well as phospholipids, Factor Xa or the von Willebrand factor binding site. However, the specific epitopes according to the invention or their major parts are preferred selected portions of said binding sites or may include a possible overlapping with said binding sites.

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In other cases, the epitopes according to the invention are more specific portions of known epitopes. Therefore, an artificial epitope could be easily obtained by synthesis and the specific above-described fragments can be deleted from non-epitopic portions such as the fragment described in a C2 fragment (amino acids phenylalanine 2196 to tryptophan 2203 inclusive and amino acids valine 2222 to phenylalanine 2234 inclusive, or the sequence leucine 2252 to threonine 2272 inclusive or the amino acids phenylalanine 2290 to threonine 2291 inclusive as well as the amino acids leucine 2306 to methionine 2321 inclusive).

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These sequences, these fragments and these epitopes are particularly advantageously characterized by high hydrophilicity, which has been defined by Parker and Hodges (1986), considerable flexibility, which has been defined by Karplus and Schultz (1985) and considerable accessibility, which has been defined by Janin (1979).

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These fragments and these epitopes are, in particular, exposed on the surface of the factor VIII protein and exhibit pronounced antigenic and immunogenic characteristics.

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Another aspect of the present invention is related to a modified (recombinant or transgenic) FVIII, possibly obtained by genetic engineering, and deleted from one or



more of the above-identified fragments, epitopes or major parts of said epitopes and/or said fragments.

Advantageously, said FVIII still allows the binding of coagulation factor(s), but will be less immunogenic and will not induce or induce less the formation of inhibitors directed against said modified FVIII or natural FVIII.

Advantageously, said polypeptide sequences, fragments or epitopes are also independently immunogenic (that is to say they are immunogenic even without being complexed with a protein of large size such as BSA, KLH haemocyanin, etc.), and preferably exhibit an immunoaffinity within inhibitors of factor VIII, such as anti-factor VIII antibodies, and/or exhibit an immunoaffinity for the receptors of the T lymphocytes and possibly B lymphocytes.

These sequences, fragments, epitopes and/or major parts of said fragments or said epitopes induce an immune reaction (antibody synthesis) when they are injected into a rabbit.

These characteristics are particularly pronounced in the case of the epitopes SEQ ID NO.:2 and SEQ ID NO.:5, which comprise sequences which are relatively "long" in amino acids, i.e. comprise 16 and 22 amino acids, respectively.

Said sequences are unexpectedly characterized by substantial immunogenicity towards monoclonal and polyclonal antibodies, but are sufficiently short to be readily and advantageously obtained by synthesis.

The present invention also relates to the conformational epitopes which comprise at least two different fragments of said sequence, at least two sequence epitopes and/or at least two major parts of said epitopes or said different fragments according to the invention and above identified.

The conformational epitopes are made up of two or more different portions of a polypeptide sequence, which portions are located in proximity to each other when the protein is folded in its tertiary or quaternary structure.

These epitopes are capable of being "recognized" (that is to say of exhibiting an immunoaffinity), preferably simultaneously, with inhibitors of factor VIII, in particular B and T lymphocytes (by way of the major histocompatibility locus (MHC I and/or II)) and/or anti-factor VIII antibodies (Scandella et al. (2000); Reding et al. (2000)).

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Preferably, the said sequence, said fragments, said epitopes and/or the major parts of said epitopes or said fragments are complexed with a carrier protein or a carrier peptide, such as BSA, or KLH haemocyanin, as to form a complex exhibiting a more powerful immunogenicity.

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The present invention is also related to a pool of more than three of said fragments, epitopes or major parts of said epitopes having advantageously important antigenic and/or immunogenic properties and which may be used advantageously in a diagnostic or therapeutic method or device such as a kit or a dialysis column, etc. allowing an efficient, preferably complete, screening and characterization of the major (if not all) known inhibitors directed against factor VIII by (human) patients.

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Another aspect of the present invention relates to an inhibitor of factor VIII which exhibits an immunoaffinity with antigenic polypeptide sequences according to the present invention, with fragments and epitopes of said sequences, with the major parts of said epitopes or said fragments and/or with the complex according to the invention.

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An inhibitor is understood to mean any biological molecule or cell (such as a T-lymphocyte) binding to said FVIII and capable of giving rise to immune disorders (characterized by humoral immune response and/or cellular immune response against said FVIII).

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In particular, such an inhibitor can be an anti-factor VIII monoclonal or polyclonal antibody or antibody fragment (such as the hypervariable Fab portion of the said antibody) which inactivates the said factor VIII and/or which inhibits the binding of factor VIII to the von Willebrand factor and/or to membrane phospholipids.

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Advantageously, the said inhibitors are synthesized by a "chimeric" animal which comprises a human immune system, such as an hu-SCID mouse or transgenic mouse producing human antibodies or other antibodies production technologies as phage display technology or immortalized B-cells, by EPV in particular.

Another aspect of the invention relates to an anti-inhibitor which is directed against the said previously described factor VIII inhibitor.

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An anti-inhibitor which is directed against the factor VIII inhibitor is understood to mean any chemical or biological molecule, a cell and/or a cell fragment (receptor)

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which is capable of interfering with the said inhibitor in such a way as to ensure its inactivation or avoid or reduce its binding to the factor VIII.

Preferably, such an anti-inhibitor is an anti-anti-factor VIII idiotype (monoclonal or polyclonal) antibody or antibody fragment, natural or obtained by genetic engineering.

Another aspect of the invention relates to a pharmaceutical composition which comprises an adequate pharmaceutical carrier or a diluent and an element selected from the group consisting of said antigenic polypeptide sequence of factor VIII, fragments and epitopes of this sequence or a pool thereof, an inhibitor of factor VIII which is directed against them, an anti-inhibitor which is directed against the said inhibitor, and/or a mixture of these.

The type and amount of adequate pharmaceutical carrier or diluent (and possibly adjuvant or excipient) present in said pharmaceutical composition, may vary according to the method of administration and is possibly combined an adjuvant in order to improve therapeutical properties of the pharmaceutical composition according to the invention or to reduce its possible side effects. Suitable pharmaceutical acceptable carriers used in the pharmaceutical composition according to the invention are well known by the person skilled in the art and are selected according to the methods generally applied by pharmacists and may include solid, liquid or gaseous non-toxic pharmaceutically acceptable carriers. The percentage of active product / pharmaceutical acceptable carrier may vary within very large ranges only limited by the tolerance and the possible side effects on patients (including humans), and by frequency and/or mode of administration.

Another aspect of the invention relates to a diagnostic and/or purification device, such as a diagnostic kit, an affinity filter, or a chromatography column which comprises an element which is selected from the group consisting of these antigenic polypeptide sequences, fragments and epitopes and/or major parts of said epitopes or said fragments, the complex according to the invention or a pool thereof, an inhibitor which is directed against them, an anti-inhibitor which is directed against said inhibitor, and/or a mixture of these. Advantageously, said device comprises a pool of said epitopes which allow a screening of patients and may detect the most important inhibitors present in said

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patients and which allow a positive test with enough specificity and sensibility.

The purification device can therefore consist of a chromatography column which comprises these sequences of factor VIII, fragments and epitopes and/or major parts of said fragments or epitopes, attached to the solid phase of the chromatography column.

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A physiological liquid (such as serum), which is derived from a patient and which comprises inhibitors of factor VIII pass through this chromatography column, with said inhibitors (for example antibodies) becoming attached specifically to said factor VIII sequences, fragments, epitopes or said major parts or a pool thereof. Following elution, it is possible to collect said inhibitors by causing them to react with anti-inhibitors (anti-anti-factor VIII idiotype antibodies).

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It is also possible to characterize the anti-anti-factor VIII idiotype antibodies which are present in a serum by these anti-inhibitors passed through a chromatography column on which inhibitors of factor VIII have been attached to the solid phase.

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It is also possible to reinject (ex vivo treatment) the physiological liquid (blood or serum or a derived fraction) to said patient after its inhibitors of factor VIII have been removed by binding with said factor VIII fragments, epitopes or a pool thereof; said inhibitors being removed from the physiological fluid (blood or serum) similarly as proposed for dialysis method applied to human patients.

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The present invention is also related to a method of treatment (ex vivo treatment) of a patient suffering from a pathology induced by inhibitors to the factor VIII which comprises the steps of extracting said physiological liquid (blood or serum) from the patient, obtaining its reaction upon a solid support binding the factor VIII fragments, epitopes or a pool thereof according to the invention and reinjecting said physiological liquid to the patient after the removing of the inhibitors having fixed said factor VIII fragments, epitopes, majors parts or a pool thereof.

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A final aspect of the invention relates to the use of the pharmaceutical composition according to the invention for preparing a medicament used for preventing and/or treating immune disorders, in particular those induced by inhibitors of factor VIII, inhibitors of the binding of factor VIII to the factor IX and/or the factor X and/or the von Willebrand factor (vWF) and/or inhibitors of the binding of factor VIII to membrane phospholipids.

The present invention will be described in details in the following non-limiting examples in reference to the enclosed figures.

Examples

Materials and Methods

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Reagents: MAS530p (Harlan-Seralab, Indianapolis, IN) is a mouse monoclonal antibody specific for the 44-kDa A2 domain of the factor VIII heavy chain. Biotin-labeled rabbit IgG anti-mouse IgG was purchased from Dakopatts (Copenhagen, Denmark). Biotin-labeled goat IgG anti-human IgG and biotin-labeled mouse IgG antirabbit IgG were obtained from Sigma Chemicals (St Louis, MI), purified α-thrombin (3000 IU/mg), streptavidin-peroxidase conjugate, ovalbumin (OVA), bovine serum albumin (BSA), keyhole limpet haemocyanin (KLH), and o-phenylenediamine (OPD) were purchased from Sigma Chemicals (St. Louis, MI). Casein was obtained from Merck (Darmstadt, Germany). 4-chloro-1-naphtol and biotinylated molecular weight markers were obtained from Bio-Rad Laboratories (Hercules, CA). Freund's adjuvant was from Difco (Detroit, Michigan).

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FVIII concentrates: Plasma FVIII (p-FVIII) was a solvent/detergent-treated FVIII concentrate (100 IU/mg protein) purified by ion exchange chromatography (FVIII Conc. SD, CAF-DCF- Red Cross, Brussels, Belgium). Albumin-free recombinant FVIII (rFVIII) was obtained from Hyland (Glendale, CA).

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<u>Plasma fraction immunoglobulins:</u> Cohn Fraction II+III was obtained from large plasma pool from 4,800 unpaid donors, after precipitation in the presence of increasing ethanol concentration. This fraction contains all Ig classes and subclasses. IgG composition was determined by nephelometry. The relative percentage of each subclass was 63,7; 30,1; 3,4 and 2,8 for IgG1, IgG2, IgG3 and IgG4 respectively (average values for 3 different batches of FII+III).

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Factor VIII concentrates, Factor VIII activity and activity inhibition

Factor VIII activity was determined in a one-stage clotting assay adapted for use on the Coagulometer KC4A (Sigma Diagnostics). The assay uses severe hemophilia A plasma (Organon Teknika, Cambridge, UK) and APPT reagent from Instrumentation Laboratory (Warrington, UK). Potencies were calculated relative to the 5th International Standard FVIII concentrate 88/640 (5.4 IU/ml) (NIBSC, Potters Bar, UK). FVIII-

inhibitory activity was measured in purified rabbit and human IgG preparations according to the modified Bethesda assay. Briefly, affinity-purified IgGs were serially diluted and incubated for 1 h in the presence of FVIII concentrate 88/640 (1 IU/ml) at 37°C. The residual FVIII activity was measured as described above.

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Activation of factor VIII by α -thrombin and immunoblotting has been described elsewhere (Peerlinck et al, 1997).

Synthesis of peptides, conjugation of peptides to carrier proteins and production of rabbit anti-peptide antisera were performed by Neosystem (Strasbourg, France).

Purification of rabbit and human antibodies by affinity chromatography

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For purification of rabbit and human antibodies, 5 mg of each different peptide was coupled to 1 ml pre-packed NHS-activated Sepharose (Pharmacia Biotech, Uppsala, Sweden) according to the manufacturer's instructions. Specific anti-peptide antibodies were purified with an automated liquid chromatography system (ÄKTAexplorer 100A, Pharmacia Uppsala, Sweden) either from 50 ml rabbit antiserum or from 100 ml of a human plasma fraction, obtained after Cohn fractionation (fraction II+III; 13 mg protein/ml). Briefly, samples were dialyzed 3 times against 5 volumes of TE buffer (20 mM Tris-HCl pH 7.2, 150 mM NaCl and 0.02% NaN₃) and loaded onto the column at a flow rate of 1 ml/min. The column was sequentially washed at 2 ml/min with 50 ml TE buffer and 30 ml TE containing 1 M NaCl. After absorption, the material was eluted (1 ml/min) with 5 ml of 0.1 M citric acid pH 2.5 and directly recovered in 5 ml of 1M Tris-HCl, pH 9.0. Samples were finally dialyzed versus 10 volumes of equilibration buffer and concentrated on Centriprep-30 (Amicon, Beverly, MA). Ig recovery was determined by the Bio-Rad protein assay.

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Selection of potential factor-VIII linear epitopes

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More than 30 surface regions (linear epitopes) spanning 8 to 25 residues, characterized by a high hydrophilicity, flexibility and accessibility were identified on the FVIII molecule. On the basis of their high probability of an outer location (see Fig. 1 for A3), 16 linear peptides (P1 to P16) were selected, matching identified stretches of 13 or more amino-acid residues. These peptides were synthesized and coupled to ovalbumin for production of specific antiserum (Table 1, hereafter). P8 includes the epitope described by Shima et al (1988) and was used as an external control.

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Experimental results obtained from said synthesized linear epitopes using the rabbit model

Results are summarized in Table 1 which concerns the characterization of rabbit anti-FVIII-peptide antisera and recovered affinity-purified of immunoglobulins.

Sixteen synthetic peptides (from 10 to 20 amino acids) were selected in the A, B, C1 and C2 domains. After conjugation with ovalbumin, the OVA-peptide conjugates were injected into rabbits and FVIII anti-peptide antisera RAP1 to RAP16 were studied.

More precisely, two rabbits were immunized with each FVIII-peptide-ovalbumin preparation. Specific antisera RAP1 to RAP16 (column b, Table 1) were prepared and assayed in an ELISA (column c, Table 1) using rFVIII or FVIII-peptide-KLH as the antigen. ELISA titer is expressed as the negative log of the reciprocal of the serum dilution giving 50% binding. The immunoglobulins were then purified by chromatography on peptide-bound Sepharose. The FVIII domain recognized by the anti-FVIII peptide Ig after immunoblotting is shown in (column d, Table 1) and Ig protein recoveries (column e, Table 1) were measured using immunoglobulins as the standard. The inhibitory activity, expressed in BU/mg protein, was determined in a FVIII neutralizing activity assay (column f, Table 1).

Immunogenicity of FVIII peptides and characterization of rabbit anti-FVIII peptide antisera

The reactivity of FVIII anti-peptide antisera was measured by an ELISA using, as antigen, either the different corresponding FVIII-peptide coupled to KLH protein or purified rFVIII. The binding reaction of each anti-FVIII-peptide antiserum was specific both for the FVIII peptide used to elicit the immune response in rabbit and for rFVIII (see Table 1).

To demonstrate the FVIII epitope specificity of the rabbit anti-peptide antibodies, rFVIII and the rFVIII fragments obtained after treatment with thrombin were resolved by SDS-PAGE and analyzed by western blotting with the different preparations of rabbit IgGs. As expected, most antisera (14/16, 87%), showed a strong

reaction with the corresponding FVIII fragment containing the selected linear epitope (see Table 1).

Purification of rabbit-anti-FVIII peptide antibodies

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The specific rabbit IgG were purified by affinity chromatography on peptide-Sepharose as described under Methods. When FVIII-neutralizing activity was measured in a one-stage clotting assay, significant inhibition was found with two rabbit IgG purified preparations: RAP2, corresponding to IgG specific for SEQ ID No. 14 and RAP7 specific for SEQ ID NO.: 1.

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Epitope mapping of rabbit anti-FVIII peptide antibodies by immunoblotting with human rFVIII

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To demonstrate the FVIII epitope specificity of the rabbit anti-peptide antibodies, rFVIII and the rFVIII fragments obtained after treatment with thrombin were resolved by SDS-PAGE and analyzed by western blotting with different preparations of rabbit IgGs (RAP1 to RAP17 Igs).

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In each run, the rFVIII heavy chain (HC) and light chain (LC) and their thrombin proteolysis products (44 kDa and 72 kDa) were identified with a mixture of two monoclonal antibodies, MoAb 530p and MoAb18, respectively specific for the heavy and light chain. MoAb18 recognizes the NH₂-terminal light-chain FVIII fragment obtained after thrombin activation, which proved too small to remain in the gel after electrophoresis. Fourteen of the 17 rabbit immunoglobulin preparations reacted strongly with both rFVIII and pFVIII. Antisera RAP1, RAP2, RAP3, RAP4 recognized exclusively the heavy chains (200 kDa to 92 kDa). Antisera RAP1 and RAP2 reacted with the 50-kDa A1-domain fragment; RAP3 and RAP4 bound to the 44-kDa fragment (domain A2); RAP5 (specific for the B domain) bound to the high-molecular-weight FVIII heavy chain (about 200-kDa).

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RAP7, RAP8, and RAP9 reacted with the 80-kDa light-chain doublet. RAP9 and RAP12 to RAP17 antibodies also detected the 72-kDa FVIII light-chain fragment. As expected, each reactive antiserum showed a strong reaction with the corresponding

FVIII fragment containing the selected linear epitope. No reaction was detectable in the gels between RAP6 or RAP10 and the HC or LC FVIII fragments.

Experimental results obtained from said synthesized linear epitopes to purify and characterize human autoantibodies

Table 2 concerns the characterization of human anti-FVIII antibodies from Cohn fraction II+III of healthy individuals.

Human anti-peptide IgG preparations (HAP1 through HAP17) were so far purified on Sepharose coupled to 13 different FVIII peptides (column a, Table 2). The Igs (column b, Table 2) were analyzed by immunoblotting. Binding to the rFVIII HC or LC chains and to the rFVIII thrombin fragment is shown respectively in columns c and d, Table 2. FVIII-domain reactivity is shown in column e, Table 2. Arrows indicate a decrease in 80-kDa band intensity. Ig recovery (column f, Table 2) after affinity purification is expressed in μg/10 mg loaded FII+III (see Materials and Methods). Inhibition of the clotting assay was determined after incubation in the presence of each of the 13 Ig preparations in the Bethesda assay (column g, Table 2).

Use of FVIII peptides for the immunopurification of human anti-FVIII antibodies in healthy donors

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To prepare and characterize human anti-FVIII antibodies present in healthy individuals, we analyzed Cohn fraction II+III, rich in immunoglobulins, for the presence of selected specific anti-peptide antibodies. Human anti-FVIII-peptide antibodies (HAP1 to HAP11, HAP16 and HAP17) were purified by affinity chromatography on Sepharose coupled to the appropriate peptide (see Table 2). As a typical example, figure 2 shows the chromatographic profile obtained with SEQ ID 32, a sequence found in C2 domain. Table 2 summarizes the results obtained with 17 epitopic sequences selected in each FVIII domain (A1, A2, A3, B, C1 and C2). Significant amounts of immunoglobulins, specific for each of the 13 FVIII peptides used, were obtained from the starting plasma fraction II+III. The specificity of the resulting purified human antibodies was directly tested by immunoblotting with plasma FVIII, recombinant FVIII, and the fragments obtained after thrombin proteolysis (see Table 2).

The IgG isotype distribution in the human purified antibody preparations was found to be quite heterogeneous. Interestingly, 40 to 79% of the recovered IgGs belonged to the IgG2 subclass. In most preparations, IgG4 appeared to be over-represented (up to 25%).

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All the human anti-FVIII-peptide antibody preparations were tested for the capacity to inhibit FVIII activity in a one-stage clotting assay. Table 2 shows that seven out of 13 preparations tested (54%) displayed inhibitory activity, SEQ ID NO.: 14, SEQ ID NO.: 19, SEQ ID NO.: 2, SEQ ID NO.: 5, SEQ ID NO.: 22, SEQ ID NO.: 32 and SEQ ID NO.: 33, respectively. As a typical example, the inhibition of FVIII activity in function of anti-SEQ ID 32 Ig concentration is shown in figure 2.

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Human anti-FVIII-peptide Ig immunospecificity towards FVIII

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The specificity of the resulting purified human antibodies was tested by immunoblotting with plasma FVIII, recombinant FVIII, and the fragments obtained after thrombin proteolysis. Again, the FVIII fragments were identified with either FVIII-HC- or FVIII-LC-specific mouse monoclonal antibodies or FVIII-peptidespecific rabbit polyclonal antibodies. The human antibodies were identified after binding of biotinylated goat anti-human IgG. Figure 3 shows the immunoreaction of high-molecular-weight FVIII (≥ 92-kDa) with four human antibody preparations, purified on Sepharose coupled to FVIII peptide SEO ID NO.: 11 (Ser¹⁰⁹- Lys¹²⁷), SEO ID NO.: 14 (Cys³²⁹- Asp³⁴⁸), SEQ ID NO.: 15 (Tyr⁴⁰⁷-Lys⁴²⁵) or SEQ ID NO.: 19 (Cys⁷¹¹-Asp⁷²⁵). The 50-kDa FVIII fragment (domain A1) was recognized by human antibodies purified on Ser¹⁰⁹- Lys¹²⁷ or Cys³²⁹-Asp³⁴⁸-Sepharose and the 44-kDa FVIII fragment (A2) by immunoglobulins purified on Tyr⁴⁰⁷-Lys⁴²⁵ and Cys⁷¹¹-Asp⁷²⁵-Sepharose. The lack of reactivity of the anti-(Ser817-Ser830) immunoglobulin preparation (HAP5) with the FVIII fragments confirms that this epitope is located in the amino-terminal end of domain B (Figure 3). Human antibodies purified on Sepharose coupled to peptide SEQ ID NO.: 1 (Arg1652-Tyr1664) or SEQ ID NO.: 2 (Asp1681-Arg1696) reacted strongly with the 80-kDa FVIII light chain (Figure 3). For both preparations, the reaction with the 80-kDa band disappeared after thrombin proteolysis, indicating that the epitopes, as expected, are located in the a3 acidic peptide at the NH₂-terminal part of the FVIII A3 domain.

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When human antibodies specific for peptide SEQ ID NO.: 5 (Arg¹⁷⁹⁷-Tyr¹⁸¹⁵ in A3 domain) were analyzed by immunoblotting, their specificity for rFVIII appeared restricted to the 80-kDa FVIII light chain and its 72-kDa thrombin fragment.

No immunoreaction with the rFVIII chains or fragments was detected with antibody preparations specific for FVIII peptides SEQ C and SEQ ID NO.: 23, although a positive reaction was obtained in the ELISA using rFVIII. This could mean that these immunoglobulin preparations recognize a conformational epitope.

Use of FVIII synthetic peptides to characterize human anti-FVIII antibodies in hemophilia A patient plasmas

The selected peptides were used in ELISA experiment to determine the anti-FVIII antibody specificity's present in hemophilia A plasmas. The peptides were coated on microplate (25 µg/ml in PBS buffer during 16h at 4°C). A 1/10 to 1/1000 dilution of plasma patient in Tris-casein buffer was reacted with the coated peptide for 2h at 37°C. The bound human IgG was measured as described in Methods. Control samples were plasma pools of healthy donors. Figure 4 shows the results obtained with the plasma of 4 hemophilia A patients. The optical densities are corrected average values (OD patient-OD normal plasma pool) of two independent experiments.

20 <u>Molecular model epitope prediction</u>

Pemberton et al (1997) have built a molecular model of the A domains of FVIII. This 3-D model makes it possible to explore predictions for important regions of FVIII activity. The model was used to locate the FVIII-peptide epitopes identified by the Parker and Hodge algorithms. As predicted by these algorithms, all peptides located in the A domains were found on the FVIII surface and were fully accessible to specific.

The overlap between the epitope and the FIXa-binding loop (5 common residues spanning Glu¹⁸¹¹-Tyr¹⁸¹⁵) may explain the inhibitory action of the corresponding anti-(Arg¹⁷⁹⁷-Tyr¹⁸¹⁵) antibodies on formation of the fibrin clot.

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<u>Analysis</u>

In the clotting test, significant inhibition of FVIII activity was recorded in the presence of rabbit anti-(Cys³²⁹-Asp³⁴⁸) and anti-(A/g¹⁶⁵³-Tyr¹⁶⁶⁴) antibodies, but different inhibition patterns were observed. Inhibition by anti-(Arg1653-Tyr1664) follows secondorder kinetics with a drastic reduction in FVI/I activity. Anti-(Cys³²⁹-Asp³⁴⁸) Ig is less efficient and shows a more complex type of reaction, with a non-linear dependence on the antibody concentration. Epitope Arg/652-Tyr¹⁶⁶⁴ and the adjacent major binding site vWF (residues Glu¹⁶⁷⁵-Glu¹⁶⁸⁴) are located in the acidic light-chain peptide a3. As shown by western blotting, a3 is released from the A3 domain after thrombin treatment. preventing further binding of anti-(Arg¹⁶⁵²-Tyr¹⁶⁶⁴) Ig to activated FVIII. Similar results have been reported by Shima et al (1991), who described the FVIII sequence Asp¹⁶⁶³-Ser¹⁶⁶⁹ as a binding site of rabbit polyclonal antibodies neutralizing FVIII activity. Epitope Cys³²⁹-Asp³⁴⁸ overlapped the acidic Asp³⁴⁸-Lys ³⁶² sequence (in a1) described as adjacent to the activated protein C (Arg³³⁶) and thrombin (Arg³⁷²) cleavage sites. It is the target of human hemophilic inhibitors. Anti-(Asp³⁴⁸-Lys³⁶²) antibodies may interfere with proteolysis or/with the FX interaction site (Met³³⁷-Arg³⁷²) (Saenko et al., 1999 and Scandella et al., 2000).

FVIII-neutralizing activity was measured in all 13 Ig preparations. Seven human Ig preparations displayed inhibition of procoagulant activity, these being specific for amino-acid residues Cys⁷¹¹-Asp⁷²⁵, Tyr¹⁶⁸¹-Arg¹⁶⁹⁶, and Arg¹⁷⁹⁷-Tyr¹⁸¹⁵ respectively. The Cys⁷¹¹-Asp⁷²⁵ sequence contains sulfated tyrosines at Tyr⁷¹⁸, Tyr⁷¹⁹, and Tyr⁷²³, and overlaps with the FVIII HC region Lys⁷¹³-Arg⁷⁴⁰ described as promoting both activation and HC proteolysis. The additional sulfated groups may be required for proper interaction with thrombin or another component as in the FX-activating complex. The sequence also overlaps with region Gly⁷⁰¹-Ser⁷⁵⁰, recognized by a weakly inhibitory mouse monoclonal antibody. Peptide P8 (Tyr¹⁶⁸¹-Arg¹⁶⁹⁶) (FVIII LC) includes the sequence Glu¹⁶⁸⁴-Arg¹⁶⁸⁹ already described by Shima et al, 1991. It contains the thrombin activation site Arg¹⁶⁸⁹-Ser¹⁶⁹⁰. P4 (Cys⁷¹¹-Asp⁷²⁵) is also included in the Asp⁷¹²-Ala⁷³⁶ sequence detected by analysis of the patient antibody repertoire by gene phage display technology. It is proposed as a possible additional inhibitor in patients (van den

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Brink et al, 2000). Peptide P9 (Arg¹⁷⁹⁷-Tyr¹⁸¹⁵) contains the FXa binding site (see below).

Of the 16 anti-FVIII-peptide immunoglobulins purified from humans or produced in rabbits, 7 did neutralize FVIII activity under the tested conditions. Using small peptide sequences and immunobinding assays, we have provided evidence for additional new epitopes. We have located new epitopes in the A1 domain (residues Ser¹⁰⁹-Cys¹²⁷), the A2 domain (Cys⁴⁰⁷-Lys⁴²⁵), and the B domain (Ser⁸¹⁷-Ser⁸³⁰ and Glu¹⁰⁷⁸-Pro¹⁰⁹²).

Autoantibodies immunopurified with denatured FVIII have been reported in healthy subjects and in pools of normal human immunoglobulins (processed fraction II, see above) (Algiman et al., 1992 and Moreau et al., 2000). A possible role in clearance of denatured FVIII or its fragments from the bloodstream and/or in the immunotolerance was suggested.

Identification of the FVIII epitopes is a major challenge to be met in order to improve FVIII treatment and the quality of therapeutic FVIII concentrates. FVIII epitope sequences help to determine the contribution of patient polyclonal anti-FVIII Igs to overall inhibitory and regulatory activity. They could also be used to monitor the usual switch in anti-FVIII specificity in a patient during treatment. Said characterization of FVIII epitopes and a model of their locations on the folded molecule improves the treatment of inhibitors in both hemophilic and non-hemophilic patients (detection, follow-up, therapeutic use of FVIII epitope peptides...).

Sub Qq

Table 1. Characterization of rabbit anti-FVIII peptides antibodies

SEQ ID(a)	Rabbit	ELISA		FVIII	RAP-IgG	Inhibitor
	Antiserum(b)	Tite	Titer(c)	domain	Recovery(e)	Titer(f)
				recognize(d)	µg/ml serum	BU/mg
		Р-КІН	r-FVIII			
SEQ ID 11	RAP1	2.5	2.2	A1	27	ı
SEQ ID 14	RAP2	3.6	2.5	A1/a1	55	1,5
SEQ ID 15 SEQ	RAP3	2.5	3.2	A2	268	,
ID 19 SEQ ID 21	RAP4	2.5	1.3	A2/a2	12	1
SEQ C	RAP5	4.6	3.9	В	106	,
SEQ ID 01 SEQ	RAP6	3.8	2.9	•	14	,
ID 02 SEQ ID 05	RAP7	3.9	3.9	a3 ↓	35	0,5
SEQ ID 23 SEQ	RAP8	1.9	6.0	a3/A3 ↓	33	•
ID 22	RAP9	3.8	2.6	A3	42	ı
SEQ ID 26 SEQ	RAP10	3.9	8.0	•	65	,
ID 27 SEQ ID 28	RAP11	ND	ND	ND	QN	QN
SEQ ID 31 SEQ	RAP12	4.1	1,1	C2	QN	ND

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ND	ND	ND	ND	ND				
ND	QN	ND	ND	ND				
C2	C2	C2	C2	C2				
1,1	6,0	0,7	1,8	1,2				
3.7	3.8	3.2	3.5	4.8				
RAP13	RAP14	RAP15	RAP16	RAP17		÷		
ID 32 SEQ ID 33						<u>.</u>		

Sub Cont

Table 2. Characterization of human anti-FVIII peptides autoantibodies

SEQ ID(a)		- FVIII	FVIII reactivity	FVIII	HAP-IgG	FVIII
		on im	on immunoblot	domain(e)	Recovery(f)	inhibitory
		-thrombin)(c) (+thrombin)(d)	rombin)(d)		µg/10 mglgG	Activity(g)
		I				BU/mg
	Anti-peptide Ig(b)	l				
SEQ ID 11	HAPI	>92kDa	50kDa	A1	0,27	1
SEQ ID 14	HAP2	>92kDa	50kDa	A1/a1	1,07	3,4
SEQ ID 15	HAP3	>92kDa	44kDa	A2	90,0	•
SEQ ID 19	HAP4	92kDa	44kDa	A2/a2	0,12	+
SEQ ID 21	HAP5	>100kDa	,	В	0,26	,
SEQ C	HAP6	ı	ı	•	0,03	
SEQ ID 01	HAP7	80kDa	80kDa	a3 ↓	0,20	
SEQ ID 02	HAP8	80kDa	80kDa	a3/A3 ↓	0,01	+
SEQ ID 05	HAP9	80kDa	72kDa	A3	0,08	+
SEQ ID 23	HAP10	ı	•	ı	0,11	ı

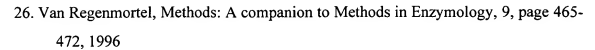
	4,3	ND	ND	ND	ND	6,3	2,4		
Sub	86,0	ND	ND	QN	ND	2,40	1,06		
V 9	QN	ND	QN	QN	QN	A3C1C2	QN		
COUT	QN	ND	ND	NO	ND	72 kDa	ND		
	QN	QN	QX	QN	QN	80kDa	ND		
	HAP11	HAP12	HAP13	HAP14	HAP15	HAP16	HAP17		
	SEQ ID 22	SEQ ID 26	SEQ ID 27	SEQ ID 28	SEQ ID 31	SEQ ID 32	SEQ ID 33		

+: Inhibition >25% at 100µg/ml

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